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(54) Title: HOMOLOGOUS 28-KILODALTON IMMUNODOMINANT PROTEIN GENES OF EHRLICHIA CANIS AND USES THEREOF

#### (57) Abstract

The present invention is directed to the cloning, sequencing and expression of homologous immnunoreactive 28-kDa protein genes, ECa28-1 and ECa28SA3, from a polymorphic multiple gene family of Ehrlichia canis. A complete sequence of another 28-kDa protein gene, ECaSA2, is also provided. Further disclosed is a multigene locus encoding all five homologous 28-kDa protein genes of Ehrlichia canis. Recombinant Ehrlichia canis 28-kDa proteins react with convalescent phase antiserum from an E. canis-infected dog.

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# HOMOLOGOUS 28-KILODALTON IMMUNODOMINANT PROTEIN GENES OF EHRLICHIA CANIS AND USES THEREOF

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#### BACKGROUND OF THE INVENTION

#### Field of the Invention

The present invention relates generally to the field of molecular biology. More specifically, the present invention relates to molecular cloning and characterization of homologous 28-kDa protein genes in *Ehrlichia canis* and a multigene locus encoding the 28-kDa homologous proteins of *Ehrlichia canis* and uses thereof.

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#### Description of the Related Art

Canine ehrlichiosis, also known as canine tropical pancytopenia, is a tick-borne rickettsial disease of dogs first described in Africa in 1935 and the United States in 1963 (Donatien and Lestoquard, 1935; Ewing, 1963). The disease became better recognized after an epizootic outbreak occurred in United States military dogs during the Vietnam War (Walker et al., 1970)

The etiologic agent of canine ehrlichiosis is Ehrlichia canis, a small, gram-negative, obligate intracellular bacterium which exhibits tropism for mononuclear phagocytes (Nyindo et al., 1971) and is transmitted by the brown dog tick, Rhipicephalus sanguineus (Groves et al., 1975). The progression of canine ehrlichiosis occurs in three phases, acute, subclinical and chronic. The acute phase is characterized by fever, anorexia, depression, lymphadenopathy and mild thrombocytopenia (Troy and Forrester, 1990). Dogs typically

recover from the acute phase, but become persistently carriers of the organism without clinical signs of disease for months or even years (Harrus et al., 1998). A chronic phase develops in some cases that characterized is by thrombocytopenia, hyperglobulinemia. anorexia, emaciation, and hemorrhage, particularly epistaxis, followed by death (Troy and Forrester, 1990).

Molecular taxonomic analysis based on the 16S rRNA gene has determined that E. canis and E. chaffeensis, the etiologic agent of human monocytic ehrlichiosis (HME), are closely related (Anderson et al., 1991; Anderson et al., 1992; Dawson et al., 1991; Chen et al., 1994). Considerable cross reactivity of the 64, 47, 40, 30, 29 and 23kDa antigens between E. canis and E. chaffeensis has been reported (Chen et al., 1994; Chen et al., 1997; Rikihisa et al., 1994; Rikihisa et Analysis of immunoreactive antigens with human and 1992). canine convalescent phase sera by immunoblot has resulted in the identification of numerous immunodominant proteins of E. canis, including a 30-kDa protein (Chen et al., 1997). In addition, a 30-kDa protein of E. canis has been described as a major immunodominant antigen recognized early in the immune response that is antigenically distinct from the 30-kDa protein of E. chaffeensis (Rikihisa et al., 1992; Rikihisa et al., 1994). Other immunodominant proteins of E canis with molecular masses ranging from 20 to 30-kDa have also been identified (Brouqui et al., 1992; Nyindo et al., 1991; Chen et al., 1994; Chen et al., 1997).

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Recently, cloning and sequencing of a multigene family (omp-1) encoding proteins of 23 to 28-kDa have been described for E chaffeensis (Ohashi et al., 1998). The 28-kDa immunodominant outer membrane protein gene (p28) of E. chaffeensis, homologous to the Cowdria ruminantium map-1 gene, was cloned. Mice immunized with recombinant P28 were protected against challenge infection with the

homologous strain according to PCR analysis of periperal blood 5 days after challenge (Ohashi et al., 1998). Molecular cloning of two similar, but nonidentical, tandemly arranged 28-kDa genes of E. canis homologous to E. chaffeensis omp-1 gene family and C. rumanintium map-1 gene has also been reported (Reddy et al., 1998).

The prior art is deficient in the lack of cloning and characterization of new homologous 28-kDa immunoreactive protein genes of *Ehrlichia canis* and a single multigene locus containing the homologous 28-kDa protein genes. Further, The prior art is deficient in the lack of recombinant proteins of such immunoreactive genes of *Ehrlichia canis*. The present invention fulfills this long-standing need and desire in the art.

#### SUMMARY OF THE INVENTION

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The present invention describes the molecular cloning, sequencing, characterization, and expression of homologous mature 28-kDa immunoreactive protein genes of Ehrlichia canis (designated Eca28-1, ECa28SA3 and ECa28SA2), and the identification of a single locus (5.592-kb) containing five 28-kDa protein genes of Ehrlichia canis (ECa28SA1, ECa28SA2, ECa28SA3, Eca28-1 and ECa28-2). Comparison with E. chaffeensis and among E. canis 28-kDa protein genes revealed that ECa28-1 shares the most amino acid homology with the E. chaffeensis omp-1 multigene family and is highly conserved among E. canis isolates. The five 28-kDa proteins were predicted to have signal peptides resulting in mature proteins, and had amino acid homology ranging from 51 to 72%. Analysis of intergenic regions revealed hypothetical promoter regions for each gene, suggesting that these genes may be independently and differentially expressed. Intergenic noncoding regions ranged in size from 299 to 355-bp. and

were 48 to 71% homologous.

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In one embodiment of the present invention, there are provided DNA sequences encoding a 30-kDa immunoreactive protein of Ehrlichia canis. Preferably, the protein has an amino acid sequence selected from the group consisting of SEQ ID No. 2, SEQ ID No. 4 and SEQ ID No. 6, and the gene has a nucleic acid sequence selected from the group consisting of SEQ ID No. 1. SEQ ID No. 3 and SEQ ID No. 5 and is a member of a polymorphic multiple gene family. Generally, the protein has an N-terminal signal sequence which is cleaved after post-translational process resulting in the production of a mature 28-kDa protein. Still preferably, the DNAs encoding 28-kDa proteins are contained in a single multigene locus, which has the size of 5.592 kb and encodes all five homologous 28-kDa proteins of Ehrlichia canis.

In another embodiment of the present invention, there is provided an expression vector comprising a gene encoding a 28-kDa immunoreactive protein of *Ehrlichia canis* and capable of expressing the gene when the vector is introduced into a cell.

In still another embodiment of the present invention, there is provided a recombinant protein comprising an amino acid sequence selected from the group consisting of SEQ ID No. 2, SEQ ID No. 4 and SEQ ID No. 6. Preferably, the amino acid sequence is encoded by a nucleic acid sequence selected from the group consisting of SEQ ID No. 1, SEQ ID No. 3 and SEQ ID No. 5. Preferably, the recombinant protein comprises four variable regions which are surface exposed, hydrophilic and antigenic. The recombinant protein may be useful as an antigen.

In yet another embodiment of the present invention, there is provided a method of producing the recombinant protein, comprising the steps of obtaining a vector that comprises an expression region comprising a sequence encoding the amino acid

sequence selected from the group consisting of SEQ ID No. 2. SEQ ID No. 4 and SEQ ID No. 6 operatively linked to a promoter; transfecting the vector into a cell; and culturing the cell under conditions effective for expression of the expression region.

The invention be may also described in embodiments as a method of inhibiting Ehrlichia canis infection in a subject comprising the steps of: identifying a subject suspected of being exposed to or infected with Ehrlichia canis; and administering a composition comprising a 28-kDa antigen of Ehrlichia canis in an amount effective to inhibit an Ehrlichia canis infection. The inhibition may occur through any means such as, i.e. the stimulation of the subject's humoral or cellular immune responses, or by other means such as inhibiting the normal function of the 28-kDa antigen, or even competing with the antigen for interaction with some agent in the subject's body.

Other and further aspects, features, and advantages of the present invention will be apparent from the following description of the presently preferred embodiments of the invention given for the purpose of disclosure.

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#### BRIEF DESCRIPTION OF THE DRAWINGS

So that the matter in which the above-recited features, advantages and objects of the invention, as well as others which will become clear, are attained and can be understood in detail, more particular descriptions of the invention briefly summarized above may be had by reference to certain embodiments thereof which are illustrated in the appended drawings. These drawings form a part of the specification. It is to be noted, however, that the appended

drawings illustrate preferred embodiments of the invention and therefore are not to be considered limiting in their scope.

Figure 1 shows nucleic acid sequence (SEQ ID No. 1) and deduced amino acid sequence (SEQ ID No. 2) of ECa28-1 gene including adjacent 5' and 3' non-coding sequences. The ATG start codon and TAA termination are shown in bold, and the 23 amino acid leader signal sequence is underlined.

Figure 2 shows SDS-PAGE of expressed 50-kDa recombinant ECa28-1-thioredoxin fusion protein (Lane 1, arrow) and 16-kDa thioredoxin control (Lane 2, arrow), and corresponding immunoblot of recombinant ECa28-1-thioredoxin fusion protein recognized by covalescent-phase *E. canis* canine antiserum (Lane 3). Thiroredoxin control was not detected by *E.canis* antiserum (not shown).

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Figure 3 shows alignment of ECa28-1 protein (SEQ ID NO. 15 2), and ECa28SA2 (partial sequence, SEQ ID NO. 7) and ECa28SA1 (SEQ ID NO. 8), E. chaffeensis P28 (SEQ ID NO. 9), E. chaffeensis OMP-1 family (SEQ ID NOs: 10-14) and C. ruminantium MAP-1 (SEQ ID NO. 15) amino acid sequences. The ECa28-1 amino acid sequence is presented as the consensus sequence. Amino acids not shown are 20 identical to ECa28-1 and are represented by a dot. Divergent amino acids are shown with the corresponding one letter abbreviation. Gaps introduced for maximal alignment of the amino acid sequences are denoted with a dash. Variable regions are underlined and denoted (VR1, VR2, VR3, and VR4). The arrows indicate the predicted signal 25 peptidase cleavage site for the signal peptide.

Figure 4 shows phylogenetic relatedness of E. canis ECa28-1 with the ECa28SA2 (partial sequence) and ECa28SA1, 6 members of the E.chaffeensis omp-1 multiple gene family, and C. rumanintium map-1 from deduced amino acid sequences utilizing unbalanced tree

construction. The length of each pair of branches represents the distance between the amino acid sequence of the pairs. The scale measures the distance between sequences.

Figure 5 shows Southern blot analysis of *E. canis* genomic DNA completely digested with six individual restriction enzymes and hybridized with a ECa28-1 DIG-labeled probe (Lanes 2-7); DIG-labeled molecular weight markers (Lanes 1 and 8).

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Figure 6 shows comparison of predicted protein characteristics of ECa28-1 (Jake strain) and E. chaffeensis P28 (Arkansas strain). Surface probability predicts the surface residues by using a window of hexapeptide. A surface residue is any residue with a >2.0 nm<sup>2</sup> of water accessible surface area. A hexapeptide with a value higher than I was considered as surface region. The antigenic index predicts potential antigenic determinants. The regions with a value above zero are potential antigenic determinants. T-cell motif locates the potential T-cell antigenic determinants by using a motif of 5 amino acids with residue 1-glycine or polar, residue 2-hydrophobic, residue 3-hydrophobic, residue 4-hydrophobic or proline, and residue 5-polar or glycine. The scale indicates amino acid positions.

Figure 7 shows nucleic acid sequences and deduced amino acid sequences of the *E. canis* 28-kDa protein genes *ECa28SA2* (nucleotide 1-849: SEQ ID No. 3; amino acid sequence: SEQ ID No. 4) and *ECa28SA3* (nucleotide 1195-2031: SEQ ID No. 5; amino acid sequence: SEQ ID No. 6) including intergenic noncoding sequences (NC2, nucleotide 850-1194: SEQ ID No. 31). The ATG start codon and termination condons are shown in bold.

Figure 8 shows schematic of the five E. canis 28-kDa protein gene locus (5.592-Kb) indicating genomic orientation and intergenic noncoding regions (28NC1-4). The 28-kDa protein genes shown in Locus 1 and 2 (shaded) have been described (McBride et al.,

1999; Reddy et al., 1998: Ohashi et al., 1998). The complete sequence of ECaSA2 and a new 28-kDa protein gene designated (ECa28SA3 - unshaded) was sequenced. The noncoding intergenic regions (28NC2-3) between ECaSA2, ECa28SA3 and ECa28-1 were completed joining the previously unlinked loci 1 and 2.

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Figure 9 shows phylogenetic relatedness of the five E canis 28-kDa protein gene members based on amino acid sequences utilizing unbalanced tree construction. The length of each pair of branches represents the distance between amino acid pairs. The scale measures the distance between sequences.

Figure 10 shows alignment of *E. canis* 28-kDa protein gene intergenic noncoding nucleic acid sequences (SEQ ID Nos. 30-33). Nucleic acids not shown, denoted with a dot (.), are identical to noncoding region 1 (28NC1). Divergence is shown with the corresponding one letter abbreviation. Gaps introduced for maximal alignment of the amino acid sequences are denoted with a dash (-). Putative transcriptional promoter regions (-10 and -35) and ribosomal binding site (RBS) are boxed.

#### DETAILED DESCRIPTION OF THE INVENTION

The present invention describes cloning, sequencing and expression of homologous genes encoding a 30-kilodalton (kDa) protein of Ehrlichia canis. A comparative molecular analysis of homologous genes among seven E. canis isolates and the E. chaffeensis omp-1 multigene family was also performed. Two new 28-kDa protein genes are identified, ECa28-1 and ECa28SA3. ECa28-1 has an 834-bp open reading frame encoding a protein of 278 amino acids (SEQ ID No. 2) with a predicted molecular mass of 30.5-kDa. An N-terminal signal sequence was identified suggesting that the protein is post-

translationally modified to a mature protein of 27.7-kDa. *ECa28SA3* has an 840-bp open reading frame encoding a 280 amino acid protein (SEQ ID No. 6).

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Using PCR to amplify 28-kDa protein genes of *E. canis*, a previously unsequenced region of *Eca28SA2* was completed. Sequence analysis of *ECa28SA2* revealed an 849-bp open reading frame encoding a 283 amino acid protein (SEQ ID No. 4). PCR amplification using primers specific for 28-kDa protein gene intergenic noncoding regions linked two previously separate loci, identifying a single locus (5.592-kb) containing all five 28-kDa protein genes. The five 28-kDa proteins were predicted to have signal peptides resulting in mature proteins, and had amino acid homology ranging from 51 to 72%. Analysis of intergenic regions revealed hypothetical promoter regions for each gene, suggesting that these genes may be independently and differentially expressed. Intergenic noncoding regions (28NC1-4) ranged in size from 299 to 355-bp, and were 48 to 71% homologous.

The present invention is directed to two new homologous 28-kDa protein genes in *Ehrlichia canis*, *Eca28-1* and *ECa28SA3*, and a complete sequence of previously partially sequenced *ECa28SA2*. Also disclosed is a multigene locus encoding all five homologous 28-kDa outer membrane proteins of *Ehrlichia canis*.

In one embodiment of the present invention, there are provided DNA sequences encoding a 30-kDa immunoreactive protein of Ehrlichia canis. Preferably, the protein has an amino acid sequence selected from the group consisting of SEQ ID No. 2, SEQ ID No. 4 and SEQ ID No. 6, and the gene has a nucleic acid sequence selected from the group consisting of SEQ ID No. 1, SEQ ID No. 3 and SEQ ID No. 5 and is a member of a polymorphic multiple gene family. More preferably, the protein has an N-terminal signal sequence which is cleaved after post-translational process resulting in the production of

a mature 28-kDa protein. Still preferably, the DNAs encoding 28-kDa proteins are contained in a single multigene locus, which has the size of 5.592 kb and encodes all five homologous, 28-kDa proteins of Ehrlichia canis.

In another embodiment of the present invention, there is provided an expression vector comprising a gene encoding a 28-kDa immunoreactive protein of *Ehrlichia canis* and capable of expressing the gene when the vector is introduced into a cell.

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In still another embodiment of the present invention, there is provided a recombinant protein comprising an amino acid sequence selected from the group consisting of SEQ ID No. 2. SEQ ID No. 4 and SEQ ID No. 6. Preferably, the amino acid sequence is encoded by a nucleic acid sequence selected from the group consisting of SEQ ID No. 1, SEQ ID No. 3 and SEQ ID No. 5. Preferably, the recombinant protein comprises four variable regions which are surface exposed, hydrophilic and antigenic. Still preferably, the recombinant protein is an antigen.

In yet another embodiment of the present invention, there is provided a method of producing the recombinant protein, comprising the steps of obtaining a vector that comprises an expression region comprising a sequence encoding the amino acid sequence selected from the group consisting of SEQ ID No. 2, SEQ ID No. 4 and SEQ ID No. 6 operatively linked to a promoter; transfecting the vector into a cell; and culturing the cell under conditions effective for expression of the expression region.

The invention may also be described in certain embodiments as a method of inhibiting Ehrlichia canis infection in a subject comprising the steps of: identifying a subject suspected of being exposed to or infected with Ehrlichia canis; and administering a composition comprising a 28-kDa antigen of Ehrlichia canis in an

amount effective to inhibit an *Ehrlichia canis* infection. The inhibition may occur through any means such as, i.e. the stimulation of the subject's humoral or cellular immune responses, or by other means such as inhibiting the normal function of the 28-kDa antigen, or even competing with the antigen for interaction with some agent in the subject's body.

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In accordance with the present invention there may be conventional molecular biology, microbiology, and employed within the skill of the art. recombinant DNA techniques Such techniques are explained fully in the literature. See, e.g., Maniatis, Fritsch & Sambrook. "Molecular Cloning: A Laboratory Manual (1982); "DNA Cloning: A Practical Approach." Volumes I and II (D.N. Glover ed. 1985); "Oligonucleotide Synthesis" (M.J. Gait ed. 1984); "Nucleic Acid Hybridization" [B.D. Hames & S.J. Higgins eds. (1985)]; "Transcription and Translation" [B.D. Hames & S.J. Higgins eds. (1984)]; "Animal Cell Culture" [R.I. Freshney, ed. (1986)]; "Immobilized Cells And Enzymes" [IRL Press, (1986)], B. Perbal, "A Practical Guide To Molecular Cloning" (1984).

Therefore, if appearing herein, the following terms shall 20 have the definitions set out below.

A "replicon" is any genetic element (e.g., plasmid, chromosome, virus) that functions as an autonomous unit of DNA replication in vivo; i.e., capable of replication under its own control.

A "vector" is a replicon, such as plasmid, phage or cosmid, to which another DNA segment may be attached so as to bring about the replication of the attached segment.

A "DNA molecule" refers to the polymeric form of deoxyribonucleotides (adenine, guanine, thymine, or cytosine) in its either single stranded form, or a double-stranded helix. This term refers only to the primary and secondary structure of the molecule.

and does not limit it to any particular tertiary forms. Thus, this term includes double-stranded DNA found, inter alia, in linear DNA molecules (e.g., restriction fragments), viruses, plasmids, and chromosomes. In discussing the structure herein according to the normal convention of giving only the sequence in the 5' to 3' direction along the nontranscribed strand of DNA (i.e., the strand having a sequence homologous to the mRNA).

A DNA "coding sequence" is a double-stranded DNA sequence which is transcribed and translated into a polypeptide in vivo when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a start codon at the 5' (amino) terminus and a translation stop codon at the 3' (carboxyl) terminus. A coding sequence can include, but is not limited to, prokaryotic sequences, cDNA from eukaryotic mRNA, genomic DNA sequences from eukaryotic (e.g., mammalian) DNA, and even synthetic DNA sequences. A polyadenylation signal and transcription termination sequence will usually be located 3' to the coding sequence.

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Transcriptional and translational control sequences are

20 DNA regulatory sequences, such as promoters, enhancers,
polyadenylation signals, terminators, and the like, that provide for the
expression of a coding sequence in a host cell.

A "promoter sequence" is a DNA regulatory region capable of binding RNA polymerase in a cell and initiating transcription of a downstream (3' direction) coding sequence. For purposes of defining the present invention, the promoter sequence is bounded at its 3' terminus by the transcription initiation site and extends upstream (5' direction) to include the minimum number of bases or elements necessary initiate transcription at levels detectable background. Within the promoter sequence will be found

transcription initiation site, as well as protein binding domains (consensus sequences) responsible for the binding of RNA polymerase. Eukaryotic promoters often, but not always, contain "TATA" boxes and "CAT" boxes. Prokaryotic promoters contain Shine-Dalgarno sequences in addition to the -10 and -35 consensus sequences.

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An "expression control sequence" is a DNA sequence that controls and regulates the transcription and translation of another DNA sequence. A coding sequence is "under the control" of transcriptional and translational control sequences in a cell when RNA polymerase transcribes the coding sequence into mRNA, which is then translated into the protein encoded by the coding sequence.

A "signal sequence" can be included near the coding sequence. This sequence encodes a signal peptide, N-terminal to the polypeptide, that communicates to the host cell to direct the polypeptide to the cell surface or secrete the polypeptide into the media, and this signal peptide is clipped off by the host cell before the protein leaves the cell. Signal sequences can be found associated with a variety of proteins native to prokaryotes and eukaryotes.

The term "oligonucleotide", as used herein in referring to the probe of the present invention, is defined as a molecule comprised of two or more ribonucleotides, preferably more than three. Its exact size will depend upon many factors which, in turn, depend upon the ultimate function and use of the oligonucleotide.

The term "primer" as used herein refers to oligonucleotide, whether occurring naturally as in a purified restriction digest or produced synthetically, which is capable of acting as a point of initiation of synthesis when placed under conditions in synthesis of a primer extension product. which complementary to a nucleic acid strand, is induced, i.e., in the

presence of nucleotides and an inducing agent such as a DNA polymerase and at a suitable temperature and pH. The primer may be either single-stranded or double-stranded and must be sufficiently long to prime the synthesis of the desired extension product in the presence of the inducing agent. The exact length of the primer will depend upon many factors, including temperature, source of primer and use the method. For example, for diagnostic applications, depending on the complexity of the target sequence. oligonucleotide primer typically contains 15-25 or more nucleotides, although it may contain fewer nucleotides.

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The primers herein are selected to be "substantially" complementary to different strands of a particular target DNA This means that sequence. the primers must be sufficiently complementary to hybridize with their respective strands. Therefore. the primer sequence need not reflect the exact sequence of the For example, a non-complementary nucleotide fragment may be attached to the 5' end of the primer, with the remainder of the primer sequence being complementary to the strand. Alternatively, non-complementary bases or longer sequences can be interspersed into the primer, provided that the primer sequence has sufficient complementarity with the sequence or hybridize therewith thereby form the template for the synthesis of the extension product.

A cell has been "transformed" by exogenous or heterologous DNA when such DNA has been introduced inside the cell. The transforming DNA may or may not be integrated (covalently linked) into the genome of the cell. In prokaryotes, yeast, and mammalian cells for example, the transforming DNA may be maintained on an episomal element such as a plasmid. With respect to eukaryotic cells, a stably transformed cell is one in which the transforming DNA has become integrated into a chromosome so that

it is inherited by daughter cells through chromosome replication. This stability is demonstrated by the ability of the eukaryotic cell to establish cell lines or clones comprised of a population of daughter cells containing the transforming DNA. A "clone" is a population of cells derived from a single cell or ancestor by mitosis. A "cell line" is a clone of a primary cell that is capable of stable growth in vitro for many generations.

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Two DNA sequences are "substantially homologous" when at least about 75% (preferably at least about 80%, and most preferably at least about 90% or 95%) of the nucleotides match over the defined length of the DNA sequences. Sequences substantially homologous can be identified by comparing sequences using standard software available in sequence data banks, or in a Southern hybridization experiment under, for example, stringent conditions as defined for that particular system. Defining appropriate hybridization conditions is within the skill of the art. See, e.g., Maniatis et al., supra; DNA Cloning, Vols. I & II, supra; Nucleic Acid Hybridization, supra.

A "heterologous' region of the DNA construct is an identifiable segment of DNA within a larger DNA molecule that is not found in association with the larger molecule in nature. Thus, when the heterologous region encodes a mammalian gene, the gene will usually be flanked by DNA that does not flank the mammalian genomic DNA in the genome of the source organism. In another example, coding sequence is a construct where the coding sequence itself is not found in nature (e.g., a cDNA where the genomic coding sequence contains introns, or synthetic sequences having codons different than the native gene). Allelic variations or naturally-occurring mutational events do not give rise to a heterologous region of DNA as defined herein.

The labels most commonly employed for these studies are radioactive elements, enzymes, chemicals which fluoresce when exposed to untraviolet light, and others. A number of fluorescent materials are known and can be utilized as labels. These include, for example, fluorescein, rhodamine, auramine, Texas Red, AMCA blue and Lucifer Yellow. A particular detecting material is anti-rabbit antibody prepared in goats and conjugated with fluorescein through an isothiocyanate.

Proteins can also be labeled with a radioactive element or with an enzyme. The radioactive label can be detected by any of the currently available counting procedures. The preferred isotope may be selected from <sup>3</sup>H, <sup>14</sup>C, <sup>32</sup>P, <sup>35</sup>S, <sup>36</sup>Cl, <sup>51</sup>Cr, <sup>57</sup>Co, <sup>58</sup>Co, <sup>59</sup>Fe, <sup>90</sup>Y, <sup>125</sup>I, <sup>131</sup>I, and <sup>186</sup>Re.

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Enzyme labels are likewise useful, and can be detected by any of the presently utilized colorimetric, spectrophotometric. fluorospectrophotometric, amperometric or gasometric techniques. The enzyme is conjugated to the selected particle by reaction with molecules bridging such as carbodiimides, diisocyanates, glutaraldehyde and the like. Many enzymes which can be used in these procedures are known and can be utilized. The preferred are peroxidase, β-glucuronidase, β-D-glucosidase, β-D-galactosidase, urease, glucose oxidase plus peroxidase and alkaline phosphatase. U.S. Patent Nos. 3,654,090, 3,850,752, and 4,016,043 are referred to by way of example for their disclosure of alternate labeling material and methods.

As used herein, the term "host" is meant to include not only prokaryotes but also eukaryotes such as yeast, plant and animal cells. A recombinant DNA molecule or gene which encodes a 28-kDa immunoreactive protein of *Ehrlichia canis* of the present invention can

be used to transform a host using any of the techniques commonly known to those of ordinary skill in the art. Especially preferred is the use of a vector containing coding sequences for a gene encoding a 28-kDa immunoreactive protein of *Ehrlichia canis* of the present invention for purposes of prokaryote transformation.

Prokaryotic hosts may include E. coli, S. tymphimurium, Serratia marcescens and Bacillus subtilis. Eukaryotic hosts include yeasts such as Pichia pastoris, mammalian cells and insect cells.

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In general, expression vectors containing promoter sequences which facilitate the efficient transcription of the inserted DNA fragment are used in connection with the host. The expression vector typically contains an origin of replication, promoter(s), terminator(s), as well as specific genes which are capable of providing phenotypic selection in transformed cells. The transformed hosts can be fermented and cultured according to means known in the art to achieve optimal cell growth.

The invention includes a substantially pure DNA encoding a 28-kDa immunoreactive protein of *Ehrlichia canis*, a strand of which DNA will hybridize at high stringency to a probe containing a sequence of at least 15 consecutive nucleotides of SEQ ID No. 1 or SEQ ID No. 3 or SEQ ID No. 5. The protein encoded by the DNA of this invention may share at least 80% sequence identity (preferably 85%, more preferably 90%, and most preferably 95%) with the amino acids listed in SEQ ID No. 2 or SEQ ID No. 4 or SEQ ID No. 6. More preferably, the DNA includes the coding sequence of the nucleotides of SEQ ID No. 1 or SEQ ID No. 3 or SEQ ID No. 5, or a degenerate variant of such a sequence.

The probe to which the DNA of the invention hybridizes preferably consists of a sequence of at least 20 consecutive nucleotides, more preferably 40 nucleotides, even more preferably 50

nucleotides, and most preferably 100 nucleotides or more (up to 100%) of the coding sequence of the nucleotides listed in SEQ ID No. 1 or SEQ ID No. 3 or SEQ ID No. 5 or the complement thereof. probe useful for detecting expression the 28-kDa immunoreactive protein of Ehrlichia canis in a human cell by a method including the steps of (a) contacting mRNA obtained from the cell with the labeled hybridization probe; and (b) detecting hybridization of the probe with the mRNA.

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This invention also includes a substantially pure DNA containing a sequence of at least 15 consecutive nucleotides (preferably 20, more preferably 30, even more preferably 50, and most preferably all) of the region from the nucleotides listed in SEQ ID No 1 or SEQ ID No. 3 or SEQ ID No. 5.

By "high stringency" is meant DNA hybridization and wash conditions characterized by high temperature and low salt concentration, e.g., wash conditions of 65°C at a salt concentration of approximately 0.1 x SSC, or the functional equivalent thereof. For example, high stringency conditions may include hybridization at about 42°C in the presence of about 50% formamide; a first wash at about 65°C with about 2 x SSC containing 1% SDS; followed by a second wash at about 65°C with about 0.1 x SSC.

By "substantially pure DNA" is meant DNA that is not part of a milieu in which the DNA naturally occurs, by virtue of separation (partial or total purification) of some or all of the molecules of that milieu, or by virtue of alteration of sequences that flank the claimed DNA. The term therefore includes, for example, a recombinant DNA which is incorporated into a vector, into an autonomously replicating plasmid or virus, or into the genomic DNA of a prokaryote or eukaryote; or which exists as a separate molecule (e.g., a cDNA or a genomic or cDNA fragment produced by polymerase chain reaction

(PCR) or restriction endonuclease digestion) independent of other sequences. It also includes a recombinant DNA which is part of a hybrid gene encoding additional polypeptide sequence, e.g., a fusion protein. Also included is a recombinant DNA which includes a portion of the nucleotides listed in SEQ ID No. 1 or SEQ ID No. 3 or SEQ ID No. 5 which encodes an alternative splice variant of a gene encoding a 28-kDa immunoreactive protein of *Ehrlichia canis*.

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The DNA may have at least about 70% sequence identity to the coding sequence of the nucleotides listed in SEQ ID No. 1 or SEQ ID No. 3 or SEQ ID No. 5, preferably at least 75% (e.g. at least 80%); and most preferably at least 90%. The identity between two sequences is a direct function of the number of matching or identical positions. When a subunit position in both of the two sequences is occupied by the same monomeric subunit, e.g., if a given position is occupied by an adenine in each of two DNA molecules, then they are identical at For example, if 7 positions in a sequence position. 10 nucleotides in length are identical to the corresponding positions in a second 10-nucleotide sequence, then the two sequences have 70% sequence identity. The length of comparison sequences will generally be at least 50 nucleotides, preferably at least 60 nucleotides, more preferably at least 75 nucleotides, and' most preferably 100 nucleotides. Sequence identity is typically measured sequence analysis software (e.g., Sequence Analysis Software Package Genetics Computer Group, University of Biotechnology Center, 1710 University Avenue, Madison, WI 53705).

The present invention comprises a vector comprising a DNA sequence coding for a which encodes a gene encoding a 28-kDa immunoreactive protein of *Ehrlichia canis* and said vector is capable of replication in a host which comprises, in operable linkage: a) an origin of replication; b) a promoter; and c) a DNA sequence coding

for said protein. Preferably, the vector of the present invention contains a portion of the DNA sequence shown in SEQ ID No. 1 or SEQ ID No. 3 or SEQ ID No. 5.

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A "vector" may be defined as a replicable nucleic acid construct, e.g., a plasmid or viral nucleic acid. Vectors may be used amplify and/or express nucleic acid encoding a 28-kDa immunoreactive protein of Ehrlichia canis. An expression vector is a replicable construct in which a nucleic acid sequence encoding a polypeptide is operably linked to suitable control sequences capable of effecting expression of the polypeptide in a cell. The need for such control sequences will vary depending upon the cell selected and the transformation method chosen. Generally, control sequences include a transcriptional promoter and/or enhancer, suitable mRNA ribosomal binding sites, and sequences which control the termination transcription and translation. Methods which are well known to those skilled in the art can be used to construct expression vectors containing appropriate transcriptional and translational control signals. See for example, the techniques described in Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual (2nd Ed.), Cold Spring Harbor Press, N.Y. A gene and its transcription control sequences are defined as being "operably linked" if the transcription sequences effectively control the transcription of the gene. Vectors of the invention include, but are not limited to, plasmid vectors and viral vectors. Preferred viral vectors of the invention are those derived from retroviruses, adenovirus, adeno-associated virus, SV40 virus, or herpes viruses.

By a "substantially pure protein" is meant a protein which has been separated from at least some of those components which naturally accompany it. Typically, the protein is substantially pure when it is at least 60%, by weight, free from the proteins and other

naturally-occurring organic molecules with which it is naturally associated in vivo. Preferably, the purity of the preparation is at least 75%, more preferably at least 90%, and most preferably at least 99%, by weight. A substantially pure 28-kDa immunoreactive protein of Ehrlichia canis may be obtained, for example, by extraction from a natural source; by expression of a recombinant nucleic acid encoding a 28-kDa immunoreactive protein of Ehrlichia canis; or by chemically synthesizing the protein. Purity can be measured by any appropriate method, e.g., column chromatography such as immunoaffinity chromatography using an antibody specific for 28-kDa a immunoreactive protein of Ehrlichia canis, polyacrylamide electrophoresis, or HPLC analysis. A protein is substantially free of naturally associated components when it is separated from at least some of those contaminants which accompany it in its natural state. Thus, a protein which is chemically synthesized or produced in a cellular system different from the cell from which it naturally originates will be, by definition, substantially free from its naturally associated components. Accordingly, substantially pure proteins include eukaryotic proteins synthesized in E. coli, other prokaryotes, or any other organism in which they do not naturally occur.

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In addition to substantially full-length proteins, the invention also includes fragments (e.g., antigenic fragments) of the 28-kDa immunoreactive protein of *Ehrlichia canis* (SEQ ID No. 2 or SEQ ID No. 4 or SEQ ID No. 6). As used herein, "fragment," as applied to a polypeptide, will ordinarily be at least 10 residues, more typically at least 20 residues, and preferably at least 30 (e.g., 50) residues in length, but less than the entire, intact sequence. Fragments of the 28-kDa immunoreactive protein of *Ehrlichia canis* can be generated by methods known to those skilled in the art, e.g., by enzymatic digestion of naturally occurring or recombinant 28-kDa immunoreactive

protein of Ehrlichia canis, by recombinant DNA techniques using an expression vector that encodes a defined fragment of 28-kDa immunoreactive protein of Ehrlichia canis, or by chemical synthesis. The ability of a candidate fragment to exhibit a characteristic of 28kDa immunoreactive protein of Ehrlichia canis (e.g., binding to an antibody specific for 28-kDa immunoreactive protein of Ehrlichia canis) can be assessed by methods described herein. Purified 28-kDa immunoreactive protein of Ehrlichia canis or antigenic fragments of 28-kDa immunoreactive protein of Ehrlichia canis can be used to generate new antibodies or to test existing antibodies (e.g., as positive controls in a diagnostic assay) by employing standard protocols known to those skilled in the art. Included in this invention are antisera generated by using 28-kDa immunoreactive protein of Ehrlichia canis or a fragment of 28-kDa immunoreactive protein of Ehrlichia canis as the immunogen in, e.g., rabbits. Standard protocols for monoclonal and polyclonal antibody production known to those skilled in this art are employed. The monoclonal antibodies generated by this procedure can be screened for the ability to identify recombinant Ehrlichia canis cDNA clones, and to distinguish them from known cDNA clones.

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Further included in this invention are fragments of the 28-kDa immunoreactive protein of *Ehrlichia canis* which are encoded at least in part by portions of SEQ ID No. 1 or SEQ ID No. 3 or SEQ ID No. 5, e.g., products of alternative mRNA splicing or alternative protein processing events, or in which a section of the sequence has been deleted. The fragment, or the intact 28-kDa immunoreactive protein of *Ehrlichia canis*, may be covalently linked to another polypeptide, e.g. which acts as a label, a ligand or a means to increase antigenicity.

The phrase "pharmaceutically acceptable" refers to 30 molecular entities and compositions that do not produce an allergic

or similar untoward reaction when administered to a human. The preparation of an aqueous composition that contains a protein as an active ingredient is well understood in the art. Typically, such compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid prior to injection can also be prepared. The preparation can also be emulsified.

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A protein may be formulated into a composition in a neutral or salt form. Pharmaceutically acceptable salts, include the acid addition salts (formed with the free amino groups of the protein) and which are formed with inorganic acids such as, for example, hydrochloric or phosphoric acids, or such organic acids as acetic, oxalic, tartaric, mandelic, and the like. Salts formed with the free carboxyl groups can also be derived from inorganic bases such as, for example, sodium, potassium, ammonium, calcium, hydroxides, and such organic bases as isopropylamine, trimethylamine, histidine, procaine and the like.

Upon formulation, solutions will be administered in a manner compatible with the dosage formulation and in such amount as is therapeutically effective. The formulations are easily administered in a variety of dosage forms such as injectable solutions.

For parenteral administration in an aqueous solution, for example, the solution should be suitably buffered if necessary and the liquid diluent first rendered isotonic with sufficient saline or glucose. These particular aqueous solutions are especially suitable intravenous. intramuscular, subcutaneous and intraperitoneal administration. In this connection, sterile aqueous media which can be employed will be known to those of skill in the art in light of the present disclosure. For example, one dosage could be dissolved in 1 mL of isotonic NaCl solution and either added to 1000mL of

hypodermoclysis fluid or injected at the proposed site of infusion, (see for example, "Remington's Pharmaceutical Sciences" 15th Edition, pages 1035-1038 and 1570-1580). Some variation in dosage will necessarily occur depending on the condition of the subject being treated. The person responsible for administration will, in any event, determine the appropriate dose for the individual subject.

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As is well known in the art, a given polypeptide may vary in its immunogenicity. It is often necessary therefore to couple the immunogen (e.g., a polypeptide of the present invention) with a carrier. Exemplary and preferred carriers are keyhole limpet hemocyanin (KLH) and human serum albumin. Other carriers may include a variety of lymphokines and adjuvants such as IL2, IL4, IL8 and others.

Means for conjugating a polypeptide to a carrier protein 15 well known in the art and include glutaraldehyde, m'maleimidobenzoyl-N-hydroxysuccinimide ester. carbo-diimide and bis-biazotized benzidine. It is also understood that the peptide may be conjugated to a protein by genetic engineering techniques that are well known in the art.

As is also well known in the art, immunogenicity to a particular immunogen can be enhanced by the use of non-specific stimulators of the immune response known as adjuvants. Exemplary and preferred adjuvants include complete BCG, Detox, (RIBI, Immunochem Research Inc.) ISCOMS and aluminum hydroxide adjuvant (Superphos, Biosector).

As used herein the term "complement" is used to define the strand of nucleic acid which will hybridize to the first nucleic acid sequence to form a double stranded molecule under stringent conditions. Stringent conditions are those that allow hybridization between two nucleic acid sequences with a high degree of homology,

but precludes hybridization of random sequences. For example, hybridization at low temperature and/or high ionic strength is termed low stringency and hybridization at high temperature and/or low ionic strength is termed high stringency. The temperature and ionic strength of a desired stringency are understood to be applicable to particular probe lengths, to the length and base content of the sequences and to the presence of formamide in the hybridization mixture.

As used herein, the term "engineered" or "recombinant" cell is intended to refer to a cell into which a recombinant gene, such as a gene encoding an Ehrlichia chaffeensis antigen has been Therefore, engineered cells are distinguishable from introduced. naturally occurring cells which do not contain a recombinantly introduced gene. Engineered cells are thus cells having a gene or genes introduced through the hand of man. Recombinantly introduced genes will either be in the form of a cDNA gene, a copy of a genomic gene, or will include genes positioned adjacent to a promoter not naturally associated with the particular introduced gene. In addition, the recombinant gene may be integrated into the host genome, or it may be contained in a vector, or in a bacterial genome transfected into the host cell.

The following examples are given for the purpose of illustrating various embodiments of the invention and are not meant to limit the present invention in any fashion.

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#### EXAMPLE 1

#### Ehrlichiae and Purification

Ehrlichia canis (Florida strain and isolates Demon, DJ, 30 Jake, and Fuzzy) were provided by Dr. Edward Breitschwerdt, (College

of Veterinary Medicine, North Carolina State University, Raleigh, NC). E. canis (Louisiana strain) was provided by Dr. Richard E. Corstvet (School of Veterinary Medicine, Louisiana State University, Baton Rouge, LA) and E. canis (Oklahoma strain) was provided by Dr. Jacqueline Dawson (Centers for Disease Control and Prevention, Atlanta, GA). Propagation of ehrlichiae was performed in DH82 cells with DMEM supplemented with 10% bovine calf serum and 2 mM Lglutamine at 37°C. The intracellular growth in DH82 cells was monitored by presence of E. canis morulae using general cytologic staining methods. Cells were harvested when 100% of the cells were infected with ehrlichiae and were then pelleted in a centrifuge at 17,000 x g for 20 min. Cell pellets were disrupted with a Braun-Sonic 2000 sonicator twice at 40W for 30 sec on ice. Ehrlichiae were purified as described previously (Weiss et al., 1975). The lysate was loaded onto discontinuous gradients of 42%-36%-30% renografin, and centrifuged at 80,000 x g for 1 hr. Heavy and light bands containing ehrlichiae were collected and washed with sucrose-phosphateglutamate buffer (SPG, 218 mM sucrose, 3.8 mM KH<sub>2</sub>PO<sub>4</sub>, 7.2 mM K<sub>2</sub>HPO<sub>4</sub>, 4.9 mM glutamate, pH 7.0) and pelleted by centrifugation.

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#### **EXAMPLE 2**

### Nucleic Acid Preparation

Ehrlichia canis genomic DNA was prepared resuspending the renografin-purified ehrlichiae in 600 µl of 10 mM 25 Tris-HCl buffer (pH 7.5) with 1% sodium dodecyl sulfate (SDS, w/v) and 100 ng/ml of proteinase K as described previously (McBride et al., 1996). This mixture was incubated for 1 hr at 56° C, and the nucleic acids were extracted twice with o f 30 phenol/chloroform/isoamyl alcohol (24:24:1). DNA was pelleted by

absolute ethanol precipitation, washed once with 70% ethanol, dried and resuspended in 10mM Tris (pH 7.5). Plasmid DNA was purified by using High Pure Plasmid Isolation Kit (Boehringer Mannheim, Indianapolis, IN), and PCR products were purified using a QIAquick PCR Purification Kit (Qiagen, Santa Clarita, CA).

#### **EXAMPLE 3**

#### PCR Amplification of the E. canis 28-kDa protein Genes

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Regions of the E. canis ECa28-1 gene selected for PCR amplification were chosen based on homology observed (>90%) in the consensus sequence generated from Jotun-Hein aligorithm alignment of E. chaffeensis p28 and Cowdria ruminantium map-1 genes. Forward primer 793 (5-GCAGGAGCTGTTGGTTACTC-3') (SEQ ID NO. 16) and reverse primer 1330 (5'-CCTTCCTCCAAGTTCTATGCC-3') (SEQ ID NO. 17) corresponded to nucleotides 313-332 and 823-843 ruminantium MAP-1 and 307-326 and 834-814 of E. chaffeensis P28. E. canis (a North Carolina isolate, Jake) DNA was amplified with primers 793 and 1330 with a thermal cycling profile of 95°C for 2 min, and 30 cycles of 95°C for 30 sec, 62° C for 1 min, 72°C for 2 min followed by a 72°C extension for 10 min and 4°C hold. PCR products were analyzed on 1% agarose gels. This amplified PCR product was sequenced directly with primers 793 and 1330.

Primers specific for ECa28SA2 gene designated 46f (5'-25 ATATACTTCCTACCTAATGTCTCA-3', SEQ ID No. 18) and primer 1330 (SEQ ID No. 17) were used to amplify the targeted region. amplified product was gel purified and cloned into a TA cloning vector (Invitrogen, Santa Clarita. CA). The clone was sequenced bidirectionally with primers: M13 reverse from the vector, 46f, ECa28SA2 (5'-AGTGCAGAGTCTTCGGTTTC-3', SEQ ID No. 19), ECa5.3 30

(5'-GTTACTTGCGGAGGACAT-3'. SEQ ID No. 20). DNA was amplified with a thermal cycling profile of 95°C for 2 min, and 30 cycles of 95°C for 30 sec, 48°C for 1 min, 72°C for 1 min followed by a 72°C extension for 10 min and 4°C hold.

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#### EXAMPLE 4

#### Sequencing Unknown 5' and 3' Regions of the ECa28-1 Gene

The full length sequence of ECa28-1 was determined using a Universal GenomeWalker Kit (CLONTECH, Palo Alto, CA) according to 10 Genomic E. canis (Jake the protocol supplied by the manufacturer. isolate) DNA was digested completely with five restriction enzymes (DraI, EcoRV, PvuII, ScaI, StuI) which produce blunt-ended DNA. An adapter (AP1) supplied in the kit was ligated to each end of E. canis The genomic libraries were used as templates to find the 15 unknown DNA sequence of the ECa28-1 gene by PCR using a primer complementary to a known portion of the ECa28-1 sequence and a primer specific for the adapter AP1. Primers specific for ECa28-1 used for genome walking were designed from the known DNA sequence derived from PCR amplification of ECa28-1 with primers 793 (SEQ ID 20 (SEQ ID NO. 17). **Primers** 394 and 1330 NO. 16) GCATTTCCACAGGATCATAGGTAA-3'; nucleotides 687-710, SEQ ID NO. 21) and 394C (5'-TTACCTATGATCCTGT GGAAATGC-3; nucleotides 710-687, SEQ ID NO. 22) were used in conjunction with supplied primer API to amplify the unknown 5' and 3' regions of the ECa28-1 25 gene by PCR. A PCR product corresponding to the 5' region of the ECa28-1 gene amplified with primers 394C and AP1 (2000-bp) was 793C (5'-GAGTA unidirectionally with primer sequenced ACCAACAGCTCCTGC-3', SEQ ID No. 23). A PCR product corresponding

to the 3' region of the *ECa28-1* gene amplified with primers 394 and API (580-bp) was sequenced bidirectionally with the same primers. Noncoding regions on the 5' and 3' regions adjacent to the open reading frame were sequenced, and primers EC28OM-F (5'-TCTACTTTGCACTTCC ACTATTGT-3', SEQ ID NO. 24) and EC28OM-R (5'-ATTCTTTTGCCACTATTT TTCTTT-3', SEQ ID NO. 25) complementary to these regions were designed in order to amplify the entire *ECa28-1* gene.

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#### EXAMPLE 5

#### Sequencing of E. canis isolates

DNA was sequenced with an ABI Prism 377 DNA Sequencer (Perkin- Elmer Applied Biosystems, Foster City, CA). The entire Eca28-1 genes of seven E canis isolates (four from North Carolina, and one each from Oklahoma, Florida, and Louisiana) were amplified by PCR with primers EC28OM-F (SEQ ID No. 24) and EC28OM-R (SEQ ID No. 25) with a thermal cycling profile of 95°C for 5 minutes, and 30 cycles of 95°C for 30 seconds, 62°C for 1 minutes, and 72°C for 2 minutes and a 72°C extension for 10 minutes. The resulting PCR products were bidirectionally sequenced with the same primers.

#### **EXAMPLE 6**

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#### Cloning and Expression of E. canis ECa28-1

The entire *E. canis ECa28-1* gene was PCR-amplified with primers-EC28OM-F and EC28OM-R and cloned into pCR2.1-TOPO TA cloning vector to obtain the desired set of restriction enzyme cleavage sites (Invitrogen, Carlsbad, CA). The insert was excised from pCR2.1-

TOPO with BstX I and ligated into pcDNA 3.1 eukaryotic expression vector (Invitrogen, Carlsbad, CA) designated pcDNA3.1/EC28 for subsequent studies. The pcDNA3.1/EC28 plasmid was amplified, and the gene was excised with a KpnI-XbaI double digestion and directionally ligated into pThioHis prokaryotic expression vector (Invitrogen, Carlsbad, CA). The clone (designated pThioHis/EC28) produced a recombinant thioredoxin fusion protein in Escherichia coli BL21. The recombinant fusion protein was crudely purified in the insoluble phase by centrifugation. The control thioredoxin fusion protein was purified from soluble cell lysates under native conditions using nickel-NTA spin columns (Qiagen, Santa Clarita, CA).

#### EXAMPLE 7

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### Western Immunoblot Analysis

Recombinant E. canis ECa28-1 fusion protein was subjected to SDS-polyacrylamide gel electrophoresis (SDS-PAGE) on 4-15% Tris-HCl gradient gels (Bio-Rad, Hercules, CA) and transferred to pure nitrocellulose (Schleicher & Schuell, Keene, NH) using a semi-dry transfer cell (Bio-Rad, Hercules, CA). The membrane was incubated with convalescent phase antisera from an E. canis-infected dog diluted 1:5000 for 1 hour, washed, and then incubated with an anti-canine IgG (H & L) alkaline phosphatase-conjugated affinity-purified secondary antibody at 1:1000 for 1 hour (Kirkegaard & Perry Laboratories, Gaithersburg, MD). Bound antibody was visualized with 5-bromo-4-chloro-3-indolyl phosphate/nitroblue tetrazolium (BCIP/NBT) substrate (Kirkegaard & Perry Laboratories, Gaithersburg, MD).

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#### **EXAMPLE 8**

#### Southern Blot Analysis

To determine if multiple genes homologous to the ECa28-1 5 gene were present in the E. canis genome, a genomic Southern blot analysis was performed using a standard procedure (Sambrook et al. 1989). E. canis genomic DNA digested completely with each of the restriction enzymes BanII, EcoRV, HaeII, KpnI and SpeI, which do not cut within the ECa28-1 genc, and AseI which digests ECa28-1 at 10 nucleotides 34, 43 and 656. The probe was produced by PCR amplification with primers EC28OM-F and EC28OM-R and digoxigenin (DIG)-labeled deoxynucleotide triphosphates (dNTPs) (Boehringer Mannheim, Indianapolis, IN) and digested with Asel. The digested probe (566-bp) was separated by agarose gel electrophoresis, gel-15 purified and then used for hybridization. The completely digested genomic E. canis DNA was electrophoresed and transferred to a nylon membrane (Boehringer Mannheim, Indianapolis, IN) and hybridized at 40°C for 16 hr with the ECa28-1 gene DIG-labeled probe in DIG Easy Hyb buffer according to the manufacturer's protocol (Boehringer 20 Mannheim, Indianapolis, IN). Bound probe was detected with a anti-DIG alkaline phosphatase-conjugated antibody and a luminescent substrate (Boehringer Mannheim, Indianapolis, IN) and exposed to BioMax scientific imaging film (Eastman Kodak, Rochester, NY).

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#### **EXAMPLE 9**

#### Sequence Analysis and Comparasion

E. chaffeensis p28 and C. ruminantium map-1 DNA sequences were obtained from the National Center of Biotechnology Information (NCBI) (World Wide Web site at URL:

http://www.ncbi.nlm.nih.gov/Entrez). Nucleotide and deduced amino sequences, and protein and phylogenetic analyses performed with LASERGENE software (DNASTAR, Inc., Madison, WI). Analysis of post-translational processing was performed method of McGeoch and von Heijne for signal sequence recognition using the PSORT program (McGeoch, 1985; von Heijne, 1986) (World Wide Web site at URL: PRIVATE HREF= "http://www.imcb.osakau.ac.jp/nakai/form.htm", **MACROBUTTON** HtmlResAnchor http://www.imcb.osaka-u.ac.jp/nakai/form.htm).

GenBank accession numbers for nucleic acid and amino acid sequences of the *E. canis ECa28-1* genes described in this study are: Jake, AF082744; Louisiana, AF082745; Oklahoma, AF082746; Demon, AF082747; DJ, AF082748; Fuzzy, AF082749; Florida, AF082750.

Sequence analysis of ECa28-1 from seven different strains of E. canis was performed with primers designed to amplify the entire gene. Analysis revealed the sequence of this gene was conserved among the isolates from North Carolina (four), Louisiana, Florida and Oklahoma.

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#### EXAMPLE 10

## PCR Amplification, Cloning, Sequencing and Expression of ECa28-1

Alignment of nucleic acid sequences from *E. chaffeensis*25 p28 and Cowdria ruminantium map-1 using the Jotun-Hein aligorithm produced a consensus sequence with regions of high homology (>90%). These homologous regions (nucleotides 313-332 and 823-843 of C. ruminantium map-1; 307-326 and 814-834 of E. chaffeensis p28) were targeted as primer annealing sites for PCR amplification.

30 PCR amplification of the E. canis ECa28-1 and E. chaffeensis p28 gene

was accomplished with primers 793 and 1330, resulting in a 518-bp PCR product. The nucleic acid sequence of the E. canis PCR product was obtained by sequencing the product directly with primers 793 and 1330. Analysis of the sequence revealed an open reading frame encoding a protein of 170 amino acids, and alignment of the 518-bp sequence obtained from PCR amplification of E. canis with the DNA sequence of E. chaffeensis p28 gene revealed a similarity greater than 70%, indicating that the genes were homologous. Adapter PCR with primers 394 and 793C was performed to determine the 5' and 3' segments of the sequence of the entire gene. Primer 394 produced four PCR products (3-kb, 2-kb, 1-kb, and 0.8-kb), and the 0.8-bp product was sequenced bidirectionally using primers 394 and AP1. The deduced sequence overlapped with the 3' end of the 518-bp product, extending the open reading frame 12-bp to a termination codon. An additional 625-bp of non-coding sequence at the 3' end of the ECa28-1 gene was also sequenced. Primer 394C was used to amplify the 5' end of the ECa28-1 gene with supplied primer AP1. Amplification with these primers resulted in three PCR products (3.3, 3-kb, and 2-kb). The 2-kb fragment was sequenced unidirectionally with primer 793C. The sequence provided the putative start codon of the ECa28-1 gene and completed the 834-bp open reading frame encoding a protein of 278 amino acids. An additional 144-bp of readable sequence in the 5' noncoding region of the ECa28-1 gene was generated. Primers EC28OM-F and EC28OM-R were designed from complementary non-coding regions adjacent to the ECa28-1 gene.

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The PCR product amplified with these primers was sequenced directly with the same primers. The complete DNA sequence (SEQ ID NO. 1) for the *E. canis ECa28-1* gene is shown in Figure 1. The *ECa28-1* PCR fragment amplified with these primers contained the entire open reading frame and 17 additional amino

acids from the 5' non-coding primer region. The gene was directionally subcloned into pThioHis expression vector, and E. coli (BL21) were transformed with this construct. The expressed ECa28-1-thioredoxin fusion protein was insoluble. The expressed protein had an additional 114 amino acids associated with the thioredoxin, 5 amino acids for the enterokinase recognition site, and 32 amino acids from the multiple cloning site and 5' non-coding primer region at the N-terminus. Convalescent-phase antiserum from an E. canis infected dog recognized the expressed recombinant fusion protein, but did not react with the thioredoxin control (Figure 2).

#### **EXAMPLE 11**

#### Sequence Homology

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15 The nucleic acid sequence of ECa28-1 (834-bp) and the E chaffeensis omp-1 family of genes including signal sequences (ECa28-1, omp-1A, B, C, D, E, and F) were aligned using the Clustal method to examine homology between these genes (alignment Nucleic acid homology was equally conserved (68.9%) between ECa28-20 1, and E. chaffeensis p28 and omp-1F. Other putative membrane protein genes in the E. chaffeensis omp-1 family, omp-1 D (68.2%), omp-1E (66.7%), omp-1C (64.1%), Cowdria ruminantium map-1 (61.8%), E. canis 28-kDa protein 1 gene (60%) and 28-kDa protein 2 gene (partial) (59.5%) were also homologous to ECa28-1. E 25 chaffeensis omp-1B had the least nucleic acid homology (45.1%) with E.Ca28-1.

Alignment of the predicted amino acid sequences of ECa28-1 (SEQ ID NO. 2) and *E. chaffeensis* P28 revealed amino acid substitutions resulting in four variable regions (VR). Substitutions or deletions in the amino acid sequence and the locations of variable

regions of ECa28-1 and the E. chaffeensis OMP-1 family were identified Amino acid comparison including the signal peptide revealed that ECa28-1 shared the most homology with OMP-1F (68%) of the E. chaffeensis OMP-1 family, followed by E. chaffeensis P28 (65.5%)OMP-1D (62.9%), OMP-1E (65.1%), OMP-1C (62.9%), Cowdria ruminantium MAP-1 (59.4%), E. canis 28-kDa protein 1 and 28-kDa protein 2 (partial) (55.6%)(53.6%),and OMP-1B The phylogenetic relationships (43.2%).based on amino sequences show that ECa28-1 and C. ruminantium MAP-1, Echaffeensis OMP-1 proteins, and E. canis 28-kDa proteins 1 and 2 (partial) are related (Figure 4).

### EXAMPLE 12

# 15 Predicted Surface Probability and Immunoreactivity

Analysis of *E. canis* ECa28-1 using hydropathy and hydrophilicity profiles predicted surface-exposed regions on ECa28-1 (Figure 6). Eight major surface-exposed regions consisting of 3 to 9 amino acids were identified on ECa28-1 and were similar to the profile of surface-exposed regions on *E. chaffeensis* P28 (Figure 6). Five of the larger surface-exposed regions on ECa28-1 were located in the N-terminal region of the protein. Surface-exposed hydrophilic regions were found in all four of the variable regions of ECa28-1. Ten T-cell motifs were predicted in the ECa28-1 using the Rothbard-Taylor aligorithm (Rothbard and Taylor, 1988), and high antigenicity of the ECa28-1 was predicted by the Jameson-Wolf antigenicity aligorithm (Figure 6) (Jameson and Wolf, 1988). Similarities in antigenicity and T-cell motifs were observed between ECa28-1 and *E. chaffeensis* P28.

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#### EXAMPLE 13

# Detection of Homologous Genomic Copies of ECa28-1 Gene

Genomic Southern blot analysis of *E. canis* DNA completely digested independently with restriction enzymes *BanII*, *EcoRV*, *HaeII*, *KpnI*, *SpeI*, which do not have restriction endonuclease sites in the *ECa28-1* gene, and *AseI*, which has internal restriction endonuclease sites at nucleotides 34, 43 and 656, revealed the presence of at least three homologous *ECa28-1* gene copies (Figure 5). Although *ECa28-1* has internal *Ase I* internal restriction sites, the DIG-labeled probe used in the hybridization experiment targeted a region of the gene within a single DNA fragment generated by the *AseI* digestion of the gene. Digestion with *AseI* produced 3 bands (approximately 566-bp, 850 -bp, and 3-kb) that hybridized with the *ECa28-1* DNA probe indicating the presence of multiple genes homologous to *ECa28-1* in the genome. Digestion with *EcoRV* and *SpeI* produced two bands that hybridized with the *ECa28-1* gene probe.

#### **EXAMPLE 14**

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## Identification of 28-kDa Protein Gene Locus

Specific primers designated *ECaSA3-2* (5'-CTAGGATTA GGTTATAGTATAAGTT-3', SEQ ID No. 26) corresponding to regions within *ECa28SA3* and primer 793C (SEQ ID No. 23) which anneals to a region with *ECa28-1* were used to amplify the intergenic region between gene *SA3* and *ECa28-1*. The 800-bp product was sequenced with the same primers. DNA was amplified with a thermal cycling profile of 95°C for 2 min, and 30 cycles of 95°C for 30 sec, 50°C for 1 min, 72°C for 1 min followed by a 72°C extension for 10 min and 4°C hold.

### **EXAMPLE 15**

PCR Amplification of 28-kDa Protein Genes and Identification of the Multiple Gene Locus

In order to specifically amplify possible unknown genes downstream of ECa28SA2, primer 46f specific for ECa28SA2, and primer 1330 which targets a conserved region on the 3' end of ECa28-I gene were used for amplification. A 2-kb PCR product was amplified with these primers that contained 2 open reading frames. open reading frame contained the known region of gene, ECaSA2, and a previously unsequenced 3' portion of the gene. Downstream from ECaSA2 an additional non identical, but homologous 28-kDa protein gene was found, and designated ECa28SA3. The two known loci were joined by amplification with primer SA3-2 specific for the 3' end of ECa28SA3 gene was used in conjunction with a reverse primer 793C, which anneals at 5' end of ECa28-1. An 800-bp PCR product was amplified which contained the 3' end of Eca28SA3, the intergenic region between ECa28SA3 and ECa28-1 (28NC3) and the 5' end of Eca28-1, joining the previously separate loci (Figure 8). The 849-bp open reading frame of ECa28SA2 encodes a 283 amino acid protein, and ECa28SA3 has an 840-bp open reading frame encoding a 280 The intergenic noncoding region between amino acid protein. ECa28SA3 and ECa28-1 was 345-bp in length (Figures 7 and 8)

25 EXAMPLE 16

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### Nucleic and Amino Acid Homology

The nucleic and amino acid sequences of all five *E. canis*28-kDa protein genes were aligned using the Clustal method to

30 examine the homology between these genes. The nucleic acid

homology ranged from 58 to 75% and a similar amino acid homology of ranging from 67 to 72% was observed between the *E. canis* 28-kDa protein gene members (Figure 9).

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### **EXAMPLE 17**

## Transcriptional Promoter Regions

The intergenic regions between the 28-kDa protein genes were analyzed for promoter sequences by comparison with consensus *Escherichia coli* promoter regions and a promoter from *E. chaffeensis* (Yu et al., 1997; McClure, 1985).

Putative promoter sequences including RBS, -10 and -35 regions were identified in 4 intergenic sequences corresponding to genes ECa28SA2, ECa28SA3, ECa28-1, and ECa28-2 (Figure 10). The upstream noncoding region of ECa28SA1 is not known and was not analyzed.

#### EXAMPLE 18

## 20 N-Terminal Signal Sequence

The amino acid sequence analysis revealed that entire E.canis ECa28-1 has a deduced molecular mass of 30.5-kDa and the entire ECa28SA3 has a deduced molecular mass of 30.7-kDa. Both proteins have a predicted N-terminal signal peptide of 23 amino acids (MNCKKILITTALMSLMYYAPSIS, SEQ ID No. 27), which is similar to that predicted for E. chaffeensis P28 (MNYKKILITSALISLISSLPGV SFS, SEQ ID NO. 28), and the OMP-1 protein family (Yu et al., 1998; Ohashi et al., 1998b). A preferred cleavage site for signal peptidases (SIS; Ser-X-Ser) (Oliver, 1985) is found at amino acids 21, 22, and 23 of ECa28-1. An additional putative cleavage site at amino acid position 25

(MNCKKILITTALISLMYSIPSISSFS, SEQ ID NO. 29) identical to the predicted cleavage site of *E. chaffeensis* P28 (SFS) was also present, and would result in a mature ECa28-1 with a predicted molecular mass of 27.7-kDa. Signal cleavage site of the previously reported partial sequence of ECa28SA2 is predicted at amino acid 30. However, signal sequence analysis predicted that ECa28SA1 had an uncleavable signal sequence.

## Summary

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Proteins of similar molecular mass have been identified and cloned from multiple rickettsial agents including *E. canis*, *E. chaffeensis*, and *C. ruminantium* (Reddy et al., 1998; Jongejan et al., 1993; Ohashi et al., 1998). A single locus in *Ehrlichia chaffeensis* with 6 homologous p28 genes, and 2 loci in *E. canis*, each containing some homologous 28-kDa protein genes have been previously described.

The present invention demonstrated the cloning, expression and characterization of genes encoding a mature 28-kDa protein of *E. canis* that are homologous to the *omp-1* multiple gene family of *E. chaffeensis* and the *C. ruminantium map-1* gene. Two new 28-kDa protein genes were identified, *Eca28-1* and *ECa28SA3*. Another *E.canis* 28-kDa protein gene, *ECa28SA2*, partially sequenced previously (Reddy *et al.*, 1998), was sequenced completely in the present invention. Also disclosed is the identification and characterization of a single locus in *E.canis* containing all five *E.canis* 28-kDa protein genes.

The E.canis 28-kDa protein are homologous to E.chaffeensis OMP-1 family and the MAP-1 protein of C. rumanintium. The most homologous E. canis 28-kDa proteins (ECa28SA3, ECa28-1 and ECa28-2) are sequentially arranged in the locus. Homology of these proteins ranged from 67.5% to 72.3%. Divergence among these 28-kDa proteins was 27.3% to 38.6%. E. canis 28-kDa proteins

ECa28SA1 and ECa28SA2 were the least homologous with homology ranging from 50.9% to 59.4% and divergence of 53.3 to 69.9%. Differences between the genes lies primarily in the four hypervariable regions and suggests that these regions are surface exposed and subject to selective pressure by the immune system. Conservation of ECa28-1 among seven E. canis isolates has been reported (McBride et al., 1999), suggesting that E. canis may be clonal in North America. Conversely, significant diversity of p28 among E. chaffeensis isolates has been reported (Yu et al., 1998).

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All of the E. canis 28-kDa proteins appear to be post translationally processed from a 30-kD protein to a mature 28-kD Recently, a signal sequence was identified on E. chaffeensis P28 (Yu et al., 1998), and N-terminal amino acid sequencing has verified that the protein is post-translationally processed resulting in cleavage of the signal sequence to produce a mature protein (Ohashi et al., 1998). The leader sequences of OMP-1F and OMP-1E have also been proposed as leader signal peptides (Ohashi et al., 1998). sequences identified on E. chaffeensis OMP-1F, OMP-1E and P28 are homologous to the leader sequence of E. canis 28-kDa protein. Promoter sequences for the p28 genes have not been determined regions were identified experimentally, but putative promoter comparison with consensus sequences of the RBS, -10 and -35 promoter regions of E. coli and other ehrlichiae (Yu et al., 1997; McClure, 1985). Such promoter sequences would allow each gene to potentially be transcribed and translated, suggesting that these genes may be differentially expressed in the host. Persistence of infection in dogs may be related to differential expression of p28 genes resulting in antigenic changes in vivo, thus allowing the organism to evade the immune response.

The E. canis 28-kda protein genes were found to exhibit acid and amino acid sequence homology with the Echaffeensis omp-1 gene family and C. ruminantium map-1 gene. Previous studies have identified a 30-kDa protein of E. canis that reacts with convalescent phase antisera against E. chaffeensis, but was believed to be antigenically distinct (Rikihisa et al., 1994). Findings based on comparison of amino acid substitutions in four variable regions of E. canis 28-kDa proteins support this possibility. Together these findings also suggest that the amino acids responsible for the antigenic differences between E. canis and E. chaffeensis P28 are located in these variable regions and are readily accessible to the immune system. It was reported that immunoreactive peptides were located in the variable regions of the 28-kDa proteins ruminantium, E. chaffeensis and E. canis (Reddy et al., 1998). Analysis of E. canis and E. chaffeensis P28 revealed that all of the variable regions have predicted surface-exposed amino acids. A study in dogs demonstrated lack of cross protection between E. canis and E (Dawson and Ewing, 1992). chaffeensis This observation may be related to antigenic differences in the variable regions of P28 as well as in other immunologically important antigens of these chrlichial Another study found that convalescent phase human antisera from E. chaffeensis-infected patients recognized 29/28-kDa protein(s) of E. chaffeensis and also reacted with homologous proteins of E. canis (Chen et al., 1997). Homologous and crossreactive epitopes on the E canis 28-kDa protein and E. chaffeensis P28 appear to be recognized by the immune system.

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E. canis 28-kDa proteins may be important immunoprotective antigens. Several reports have demonstrated that the 30-kDa antigen of E. canis exhibits strong immunoreactivity (Rikihisa et al., 1994; Rikihisa et al., 1992). Antibodies in

convalescent phase antisera from humans and dogs have consistently reacted with proteins in this size range from E. chaffeensis and E canis, suggesting that they may be important immunoprotective antigens (Rikihisa et al., 1994; Chen et al., 1994; Chen et al., 1997). In addition, antibodies to 30, 24 and 21-kDa proteins developed early in the immune response to E. canis (Rikihisa et al., 1994; Rikihisa et al., 1992), suggesting that these proteins may be especially important in the immune responses in the acute stage of disease. Recently, a family of homologous genes encoding outer membrane proteins molecular masses of 28-kDa have been identified in E. chaffeensis, and mice immunized with recombinant E. chaffeensis P28 appeared to have developed immunity against homologous challenge (Ohashi et The P28 of E. chaffeensis has been demonstrated to be present in the outer membrane, and immunoelectron microscopy has localized the P28 on the surface on the organism, and thus suggesting that it may serve as an adhesin (Ohashi et al., 1998). It is likely that the 28-kDa proteins of E. canis identified in this study have the same location and possibly serve a similar function.

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Comparison of ECa28-1 from different strains of E. canis revealed that the gene is apparently completely conserved. Studies involving E chaffeensis have demonstrated immunologic and molecular evidence of diversity in the ECa28-1. Patients infected with E. chaffeensis have variable immunoreactivity to the 29/28-kDa proteins, suggesting that there is antigenic diversity (Chen et al., 1997). Recently molecular evidence has been generated to support antigenic diversity in the p28 gene from E chaffeensis (Yu et al., 1998). A comparison of five E. chaffeensis isolates revealed that two isolates (Sapulpa and St. Vincent) were 100% identical, but three others (Arkansas, Jax, 91HE17) were divergent by as much as 13.4% at the amino acid level. The conservation of ECa28-1 suggests that E

canis strains found in the United States may be genetically identical, and thus E. canis 28-kDa protein is an attractive vaccine candidate for canine ehrlichiosis in the United States. Further analysis of E. canis isolates outside the United States may provide information regarding the origin and evolution of E. canis. Conservation of the 28-kDa protein makes it an important potential candidate for reliable serodiagnosis of canine ehrlichiosis.

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The role of multiple homologous genes is not known at this point; however, persistence of *E.canis* infections in dogs could conceivably be related to antigenic variation due to variable expression of homologous 28-kDa protein genes, thus enabling *E canis* to evade immune surveillance. Variation of msp-3 genes in A. marginale is partially responsible for variation in the MSP-3 protein, resulting in persistent infections (Alleman et al., 1997). Studies to examine 28-kDa protein gene expression by *E. canis* in acutely and chronically infected dogs would provide insight into the role of the 28-kDa protein gene family in persistence of infection.

The following references were cited herein.

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Any patents or publications mentioned in this specification are indicative of the levels of those skilled in the art to which the invention pertains. These patents and publications are herein incorporated by reference to the same extent as if each individual publication was individually indicated to be incorporated by reference.

One skilled in the art will readily appreciate that the present invention is well adapted to carry out the objects and obtain

the ends and advantages mentioned, as well as those inherent therein. The present examples along with the methods, procedures, treatments, molecules, and specific compounds described herein are presently representative of preferred embodiments, are exemplary, and are not intended as limitations on the scope of the invention. Changes therein and other uses will occur to those skilled in the art which are encompassed within the spirit of the invention as defined by the scope of the claims.

### WHAT IS CLAIMED IS:

- 1. DNA sequences encoding a 30-kilodalton protein of Ehrlichia canis, wherein said protein is immunoreactive with anti5. Ehrlichia canis serum.
- The DNA sequences of claim 1, wherein said protein has an amino acid sequence selected from the group consisting of SEQ
   ID No. 2, SEQ ID No. 4 and SEQ ID No. 6.
  - 3. The DNA sequences of claim 2, wherein said protein has an N-terminal signal sequence.

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4. The DNA sequences of claim 3, wherein said protein is post-translationally modified to a 28-kilodalton protein.

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5. The DNA sequences of claim 1, wherein said DNA has a sequence selected from the group consisting of SEQ ID No. 1, SEQ ID No. 3 and SEQ ID No. 5.

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6. The DNA sequences of claim 1, wherein said DNA is contained in a single locus of Ehrlichia canis.

7. The DNA sequences of claim 6, wherein said locus is a multigene locus of 5.592 kb in length.

- 5 8. The DNA sequences of claim 7, wherein said locus encoding homologous 28-kilodalton proteins of Ehrlichia canis.
- 9. The DNA sequences of claim 8, wherein said homologous 28-kilodalton proteins of *Ehrlichia canis* are selected from the group consisting of ECa28SA1, ECa28SA2, ECa28SA3, ECa28-1 and ECa28-2.
  - 10. A vector comprising the DNA sequences of claim 1.
- 11. The vector of claim 10, wherein said vector is an expression vector capable of expressing a peptide or polypeptide encoded by the sequence selected from the group consisting of SEQ ID

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- No. 1, SEQ ID No. 3 and SEQ ID No. 5 when said expression vector is introduced into a cell.
- 12. A recombinant protein comprising the amino acid sequence selected from the group consisting of SEQ ID No. 2, SEQ ID No. 4 and SEQ ID No. 6.
- 13. The recombinant protein of claim 12, wherein said 30 amino acid sequence is encoded by a nucleic acid segment comprising

a sequence selected from the group consisting of SEQ ID No. 1, SEQ ID No. 3 and SEQ ID No. 5.

- 5 14. A host cell comprising the nucleic acid segment selected from the group consisting of SEQ ID No. 1, SEQ ID No. 3 and SEQ ID No. 5.
- 15. A method of producing the recombinant protein of claim 12, comprising the steps of:

obtaining a vector that comprises an expression region comprising a sequence encoding the amino acid sequence selected from the group consisting of SEQ ID No. 2, SEQ ID No. 4 and SEQ ID No. 6 operatively linked to a promoter;

transfecting said vector into a cell; and culturing said cell under conditions effective for expression of said expression region.

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- 16. An antibody immunoreactive with an amino acid sequence selected from the group consisting of SEQ ID No. 2, SEQ ID No. 4 and SEQ ID No. 6.
- 25 17. A method of inhibiting Ehrlichia canis infection in a subject comprising the steps of:

identifying a subject suspected of being exposed to or infected with Ehrlichia canis; and

administering a composition comprising a 28-kDa antigen of Ehrlichia canis in an amount effective to inhibit an Ehrlichia canis infection.

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18. The method of claim 17, wherein said 28-kDa antigen is a recombinant protein comprising an amino acid sequence selected from the group consisting of SEQ ID No. 2, SEQ ID No. 4 and SEQ ID No. 6.

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- 19. The method of claim 18, wherein said recombinant protein is encoded by a gene comprising a sequence selected from the group consisting of SEQ ID No. 1, SEQ ID No. 3 and SEQ ID No. 5.
- 15 20. The method of claim 18, wherein said recombinant protein is dispersed in a pharmaceutically acceptable carrier.

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1441	A	$\Pi$	TT	AC.	AT.	TAG	πG	ЮA	AA(	JT A	ιTΆ	TG	CC	AA	31	CCC	T	ነልር	ገልና	TT	TG	700	TA	TT	TT	<b>C</b> ል	٣٣	רא י	รกก
1501	A	٩GA	lag	AG.	AA.	AAA	CA	.CA	AC/	<b>AAC</b>	TG	GA	GT	П	TC	GG	TT	'AA	AA	CA	AG	ATT	GC	GA	.CG	GA	GC/	AA I	560
1561	C	\CI	AA	AG	GA"	IGC	AA	.GC	wG(	.AG	CC	AC	AC	AW.	ΓA	GA(	CCC	A	\GT	AC	AA7	$\mathbf{c}$							1607
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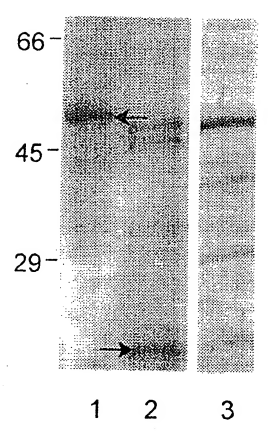
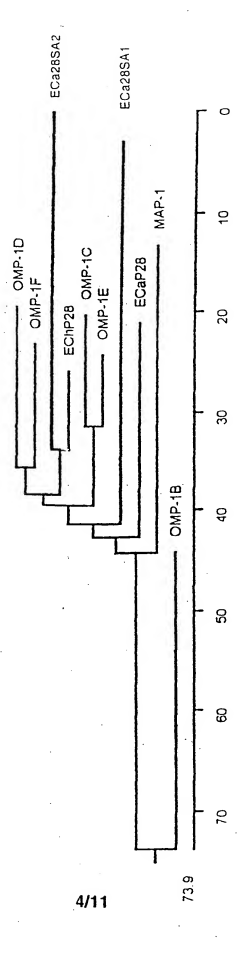


FIG. 2

<u>.</u> ධි



<u>G</u>. 4

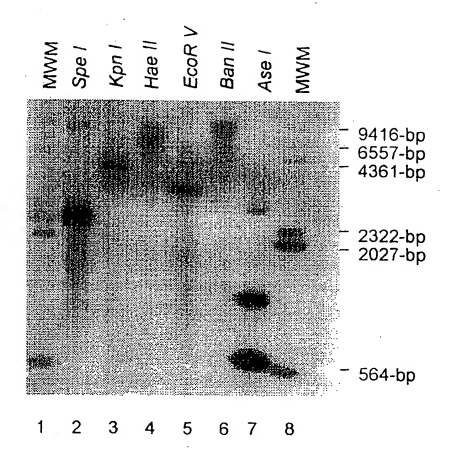
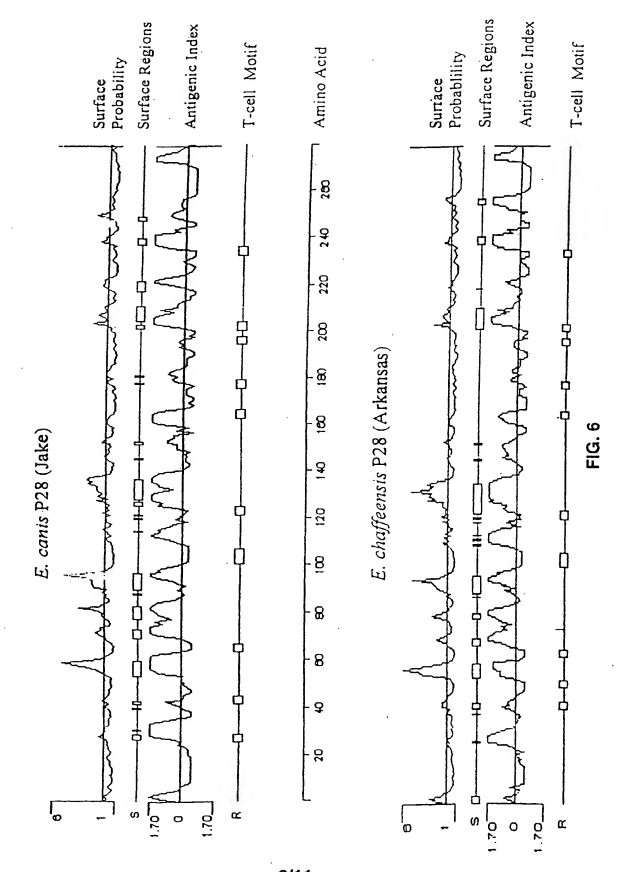


FIG. 5



#### Eca28SA2

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- AAGACAACTGTAGTATATGGCTTAAAAGAAAACTGGGCAGGAGATGCAATATCTAGTCAA 240 K T T V V Y G L K E N W A G D A I S S Q
- TTAGGGTTTGCAGTAGCTATTGGTTACTCGATAGGCAGTCCAAGAATAGAAGTTGAGATG 360 L G F A V A I G Y S I G S P R I E V E M
- TCTTATGAAGCATTTGATGTGAAAAATCCAGGTGATAATTACAAAAACGGTGCTTACAGG 420 S Y E A F D V K N P G D N Y K N G A Y R.
- TATTGTGCTTTATCTCATCAAGATGATGACGGATGATGACATGACAAA 480 Y C A L S H Q D D A D D M T S A T D K
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- CCACAGTTTGCAACAGTAACATTAAATGTATGCTACTTTGGATTAGAACTTGGATGTAGG 840

  P Q F A T V T L N V C Y F G L E L G C R

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A TOTAL COMMON TO COMMON T	
ECa28SA3 (SEQ ID NO:	31)
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COURT & & MORROW COURT OF A P. C. T. C.	1000
AGCTTCACTACTGTAGAGTGTTTATCAATGCTTTGTTTATTAATACTCTACATAATAT	1080

- ATTTGCTTTATACTTCCACTATTGTTAATTTATTTTCACTATTTTAGGTGTAATATGAAT 1200
- TGCAAAAAATTCTTATAACAACTGCATTAATGTCATTAATGTACTATGCTCCAAGCATA 1260 C K K I L I T T A L M S L M Y Y A P S I
- TCTTTTTCTGATACTATACAAGACGATAACACTGGTAGCTTCTACATCAGTGGAAAATAT 1320 S F S D T I Q D D N T G S F Y I S G K Y
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- AATATATTCACAGTTCAAAATTATTCGTTTAAATACGAAAACAACCCATTCTTAGGGTTT 1500 N I F T V Q N Y S F K Y E N N P F L G F
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- GAAGGAATGCCTTTTTCACCTTATATTTGTGCAGGTGTTGGTACTGATGTTGTTTCCATG 1800 E G M P F S P Y I C A G V G T D V V S M
- TTTGAAGCTATAAATCCTAAAATTTCTTACCAAGGAAAACTAGGATTAGGTTATAGTATA 1860 F E A I N P K I S Y Q G K L G L G Y C I
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FIG. 7-2

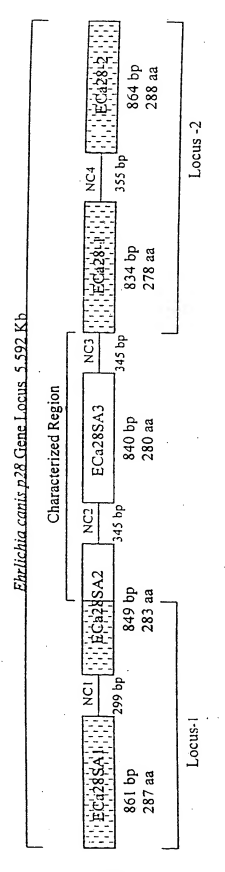
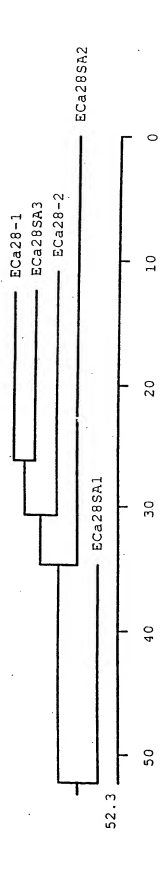


FIG. 8





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ID NO: 30) ID NO: 31) ID NO: 32) ID NO: 33)

28nc1 28nc2 28nc3 28nc3 FIG. 10

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Tyr Glu Thr Phe Asp Val Lys Asn Gln Gly Asn Asn Tyr Lys Asn
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Phe	Gly	Leu	Lys		Asp	GLY	Asp	He		GIn	Ser	Ala	Asn	
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GIÀ	Pne	ser	GTA		TTE	GIÀ	Tyr	Ala		ASP	GIY	PIO	Arg	120
<b>01.</b>	Ton	Clu	λla	110	Ф. rx-	Cln	Tvc	Dho	115	ת 1 ת	Lvc	λαη	Pro	
GIU	Leu	GIU	Ala	125	TYL	GIII	пåг	FILE	130	AIA	пуз	ASII	Pro	135
λen.	Δen	Asn	Thr		Ser	Glv	Asn	ጥኒያታ		Lvs	ጥኒያዮ	Phe	Gly	
WOII.	ASII	1150		140	001	Cly	ı iob	17.	145	275	~ y ~		011	150
Ser	Ara	Glu	Asp		Ile	Ala	Asp	Lvs		Tvr	Val	Val	Leu	
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Asn	Glu	Gly	Ile		Phe	Met	Ser			Val	Asn	Thr	Cys	
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Pro Thr Val Ala	Leu Tyr Gly Leu Lys Gln Asp Trp Asn Gly	Val
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	Gly Leu Ser Tyr Ser Ile Asn Pro Glu Ala S	
		225
	Gly His Phe His Lys Val Ala Gly Asn Glu I	
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Δla	Thr	Pro	Asn			Thr	Val	<b>.</b> 			. Vai	0.75	Hic	Phe
711.0	1111			260		1111	val	1111	265		val	Cy 5		270
Glv	Val	Glu	Leu			Ara	Phe	Asn						2,0
رين	, ,	0		275	027	9			280					
									200					
	-0	10.		1.0										
		10>		12	٠									
		11>		286										
		12>		PRT	لطمة	a ah	- F F -		_					
		13> 20>		EIILL	ICHI	a Cn	arre	ensi	S					
		23>		amin	0 30	ia c	00010	nco	of t	' ah	<i>f f c</i>	onai	~ OM	תוי מו
		00>		12	o ac	iu s	edue	nce	Or E	. CII	alie	ensi	S OM	IP-1D
Met	Asn	Cys	Glu		Phe	Phe	Ile	Thr		Ala	Leu	Thr	Leu	
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Sor	Ala	Sar	ніс		Gly	1727	Pho	Sor	40 112	Lvc	Cl.:	Cl.··	λ <b>~</b> α	45
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Thr	Thr	Val	Glv		Phe	Glv	Tle	Glu		Asp	Trp	Asp	Ara	
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Tyr	Ser	Phe	Lys	Tyr	Glu	Asn	Asn	Leu	Phe	Ser	Gly	Phe	Ala	Gly
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GIY	ASN	Giu	Pne		Asp	iie	Pro	Thr		Ile	Pro	Ser	Glu	
ח ז ת	Lou	71.	C1.,	245	C1	N ===	m	D	250				_	255
VIO	. neu	ALG	GIY	260	GIY	ASII	īŷŗ	Pro		Ile	val	Thr	Leu	
Va 1	Pho	ጥህ ፖ	Pho		Tla	Glu	T 011	C111	265	Arg	Dha	3	D1-	270
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Ser	Ser	Ser	Ser	His	Asn .	Asp A	Asn 1	His	Phe I	Asn A	Asn	Lys		
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Ser	? Phe	e Lys	з Тул	r Glu 95	Asn	Asn	Pro	) Phe	Leu 100		Phe	: Ala	Gly	
Tle	Gly	<i>r</i> ጥህነ	r Ser	Met	Glv	. Glv	Dro	720			Dho	C1.		105
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Tvr	Glu	Thr	. Phe	Asp	Val	LVS	Aen	Gln			λαη	Паст	Tira	120
-,-	0_0			125		2,3	23511	GIN	130	nsn	ASII	TYL	Lys	
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Asp	) As	n Va	1 G1		y Ası	n Phe	Y)	r Il	e Se	r Gl	y Lys	з Ту:	r Vai	l Pro
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Ser	· Va.	L Se	r Hi		e Gly	/ Val	. Phe	e Sei		a Lys	s Glr	ı Glı	ı Arç	g Asn
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Thr	Thi	r Thi	r Gly		l Ph∈	e Gly	Leu	Lys		ı Asp	Tr	Asp	Gly	ser Ser
шŗ	77.		- r	65					.70	_				75
THE	TIE	s Sei	r. r.ys		ı Şer	Pro	Glu	Asn		Phe	Asn	Val	Pro	Asn
Ma ese	Cox	Dha		80	. Cl		_		85	_			_	90
ıyı	Ser	PHE	s rys	95	GIU	Asn	Asn	Pro		Leu	Gly	Phe	Ala	Gly
ÁΊa	Va 1	Gla	т Фът		Mot	7	G1	D	100		~1			105
AIG	val	GLy	TYL	110	Mec	Asn	сту	Pro		TTE	GIu	Leu	Glu	
Ser	Tvr	Glu	Thr		Asn	Val	Lvc	) cn	115	C1	7.00	B ===		120
-	- , -	0_0		125	1150	·	ыys	ASII	130	Gry	ASII	ASI	ıyr	
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